

## Infrared Spectroscopy Absorption Table

The following table lists **infrared spectroscopy absorptions** by frequency regions.

4000-3000 cm<sup>-1</sup>

			4000-3000	cm <sup>-1</sup>			
3700-3584	medium	sharp	О-Н	stretching	alcohol	free	
3550-3200	strong	broad	О-Н	stretching	alcohol	intermolecular bonded	
3500- 3400	medium	-	N-H	stretching	primary amine	-	
3400-3300 3330-3250	medium	-	N-H	stretching	aliphatic primary amine	-	
3350-3310	medium	-	N-H	stretching	secondary amine	-	
3300-2500	strong	broad	О-Н	stretching	carboxylic acid	usually centered on 3000 cm <sup>-1</sup>	
3200-2700	weak	broad	О-Н	stretching	alcohol	intramolecular bonded	
3000-2800	strong	broad	N-H	stretching	amine salt	-	
3000-2500 cm <sup>-1</sup>							
3333-3267	strong	sharp	С-Н	stretching	alkyne	-	
2100 2000	di		CII	atuat alain w	-11		

3333-3267	strong	sharp	С-Н	stretching	alkyne	-
3100-3000	medium	-	С-Н	stretching	alkene	-
3000-2840	medium	-	С-Н	stretching	alkane	-
2830-2695	medium	-	С-Н	stretching	aldehyde	doublet
2600-2550	weak	-	S-H	stretching	thiol	-

## 2400-2000 cm<sup>-1</sup>

2349	strong	-	O=C=O	stretching	carbon dioxide	-
2275-2250	strong	broad	N=C=O	stretching	isocyanate	-
2260-2222	weak	-	CEN	stretching	nitrile	-
2260-2190	weak	-	CEC	stretching	alkyne	disubstituted
2175-2140	strong	-	S-CEN	stretching	thiocyanate	-
2160-2120	strong	-	N=N=N	stretching	azide	-
2150	-	-	C=C=O	stretching	ketene	-
2145-2120	strong	-	N=C=N	stretching	carbodiimide	-
2140-2100	weak	-	CEC	stretching	alkyne	monosubstituted
2140-1990	strong	-	N=C=S	stretching	isothiocyanate	-
2000-1900	medium	-	C=C=C	stretching	allene	-
2000	-	-	C=C=N	stretching	ketenimine	-

2000-1650 cm<sup>-1</sup>



2000-1650	weak	-	С-Н	bending	aromatic compound	overtone		
1870-1540 cm <sup>-1</sup>								
1818 1750	strong	-	C=O	stretching	anhydride	-		
1815-1785	strong	-	C=O	stretching	acid halide	-		
1800-1770	strong	-	C=O	stretching	conjugated acid halide	-		
1775 1720	strong	-	C=O	stretching	conjugated anhydride	-		
1770-1780	strong	-	C=O	stretching	vinyl / phenyl ester	-		
1760	strong	-	C=O	stretching	carboxylic acid	monomer		
1750-1735	strong	-	C=O	stretching	esters	6-membered lactone		
1750-1735	strong	-	C=O	stretching	δ-lactone	γ: 1770		
1745	strong	-	C=O	stretching	cyclopentanone	-		
1740-1720	strong	-	C=O	stretching	aldehyde	-		
1730-1715	strong	-	C=O	stretching	α,β-unsaturated ester	or formates		
1725-1705	strong	-	C=O	stretching	aliphatic ketone	or cyclohexanone or cyclopentenone		
1720-1706	strong	-	C=O	stretching	carboxylic acid	dimer		
1710-1680	strong	-	C=O	stretching	conjugated acid	dimer		
1710-1685	strong	-	C=O	stretching	conjugated aldehyde	-		
1690	strong	-	C=O	stretching	primary amide	free (associated: 1650)		
1690-1640	medium	-	C=N	stretching	imine / oxime	-		
1685-1666	strong	-	C=O	stretching	conjugated ketone	-		
1680	strong	-	C=O	stretching	secondary amide	free (associated: 1640)		
1680	strong	-	C=O	stretching	tertiary amide	free (associated: 1630)		
1650	strong	-	C=O	stretching	δ-lactam	γ: 1750-1700 β: 1760-1730		

1670-1600 cm<sup>-1</sup>



1678-1668	weak	-	C=C	stretching	alkene	disubstituted (trans)
1675-1665	weak	-	C=C	stretching	alkene	trisubstituted
1675-1665	weak	-	C=C	stretching	alkene	tetrasubstituted
1662-1626	medium	-	C=C	stretching	alkene	disubstituted (cis)
1658-1648	medium	-	C=C	stretching	alkene	vinylidene
1650-1600	medium	-	C=C	stretching	conjugated alkene	-
1650-1580	medium	-	N-H	bending	amine	-
1650-1566	medium	-	C=C	stretching	cyclic alkene	-
1648-1638	strong	-	C=C	stretching	alkene	monosubstitute
1620-1610	strong	-	C=C	stretching	α,β-unsaturated ketone	-
			1600-1300 cm <sup>-1</sup>			
1550-1500 1372-1290	strong	-	N-O	stretching	nitro compound	-
1465	medium	-	С-Н	bending	alkane	methylene gro
1450 1375	medium	-	С-Н	bending	alkane	methyl group
1390-1380	medium	-	С-Н	bending	aldehyde	-
1385-1380 1370-1365	medium	-	С-Н	bending	alkane	gem dimethyl
			1400-1000 cm <sup>-1</sup>			
1440-1395	medium	-	О-Н	bending	carboxylic acid	-
1420-1330	medium	-	О-Н	bending	alcohol	-
1415-1380 1200-1185	strong	-	S=O	stretching	sulfate	-
1410-1380 1204-1177	strong	-	S=O	stretching	sulfonyl chloride	-
1400-1000	strong	-	C-F	stretching	fluoro compound	-
1390-1310	medium	-	О-Н	bending	phenol	-
1372-1335 1195-1168	strong	-	S=O	stretching	sulfonate	-
1370-1335 1170-1155	strong	-	S=O	stretching	sulfonamide	-



1350-1342 1165-1150	strong	-	S=O	stretching	sulfonic acid	anhydrous hydrate: 1230- 1120
1350-1300 1160-1120	strong	-	S=O	stretching	sulfone	-
1342-1266	strong	-	C-N	stretching	aromatic amine	-
1310-1250	strong	-	C-O	stretching	aromatic ester	-
1275-1200 1075-1020	strong	-	C-O	stretching	alkyl aryl ether	-
1250-1020	medium	-	C-N	stretching	amine	-
1225-1200 1075-1020	strong	-	C-O	stretching	vinyl ether	-
1210-1163	strong	-	C-O	stretching	ester	-
1205-1124	strong	-	C-O	stretching	tertiary alcohol	-
1150-1085	strong	-	C-O	stretching	aliphatic ether	-
1124-1087	strong	-	C-O	stretching	secondary alcohol	-
1085-1050	strong	-	C-O	stretching	primary alcohol	-
1070-1030	strong	-	S=O	stretching	sulfoxide	-
1050-1040	strong	broad	CO-O-CO	stretching	anhydride	-
			1000-650 cm <sup>-1</sup>			
995-985 915-905	strong	-	C=C	bending	alkene	monosubstitute
980-960	strong	-	C=C	bending	alkene	disubstituted (trans)
895-885	strong	-	C=C	bending	alkene	vinylidene
850-550	strong	-	C-Cl	stretching	halo compound	-
840-790	medium	-	C=C	bending	alkene	trisubstituted
730-665	strong	-	C=C	bending	alkene	disubstituted (cis)
	strong	-	C-Br	stretching	halo compound	-
690-515	strong					
690-515 600-500	strong	-	C-I	stretching	halo compound	-
					halo compound	-
			C-I		halo compound  1,2,4- trisubstituted	-



810 ± 20	strong	-	С-Н	bending	1,4- disubstituted or 1,2,3,4- tetrasubstituted	-
$780 \pm 20$ $(700 \pm 20)$	strong	-	С-Н	bending	1,2,3- trisubstituted	-
755 ± 20	strong	-	С-Н	bending	1,2- disubstituted	-
750 ± 20 700 ± 20	strong	-	С-Н	bending	monosubstitute d benzene derivative	-

## **Contributors and Attributions**

OChemOnline

Infrared Spectroscopy Absorption Table is shared under a CC BY-NC-SA 4.0 license and was authored, remixed, and/or curated by LibreTexts.